

Liu, D. J. et al. (2017) Exome-wide association study of plasma lipids in >300,000 individuals. *Nature Genetics*, 49, pp. 1758-1766.
(doi: [10.1038/ng.3977](https://doi.org/10.1038/ng.3977))

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Deposited on: 01 November 2017

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Title: Exome-wide association study of plasma lipids in >300,000 individuals

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Plasma lipid levels are modifiable risk factors for atherosclerotic cardiovascular disease. In >300,000 participants, we screened DNA sequence variants on an exome-focused array and identified 444 independent variants in 250 loci significantly associated with total cholesterol (TC), high-density lipoprotein cholesterol (HDL-C), low-density lipoprotein cholesterol (LDL-C), and/or triglycerides (TG). At two loci (*JAK2* and *AICF*), experimental analysis in mice revealed TC and TG changes, respectively, consistent with the human data. We utilized 444 mapped variants to address several questions of clinical relevance and found the following: (1) beta-thalassemia trait carriers displayed lower total cholesterol and were protected from coronary artery disease; (2) multiple *CETP* variants each consistently associated with not only higher HDL-C but also elevated risk for age-related macular degeneration; (3) only some mechanisms of lowering LDL-C seemed to increase risk for type 2 diabetes; and (4) at TG genes involved in hepatic production of TG-rich lipoproteins (e.g., *TM6SF2*, *PNPLA3*), the plasma TG-lowering allele tracked with higher liver fat, higher risk for type 2 diabetes, and lower risk for coronary artery disease whereas at TG genes involved in peripheral lipolysis (e.g., *LPL*, *ANGPTL4*), the plasma TG-lowering allele had no effect on liver fat but lowered risks for *both* type 2 diabetes and coronary artery disease.

Main Text

Association studies testing common DNA sequence variation have uncovered 175 genetic loci that affect lipid levels in the population¹⁻⁷, with most variants residing in the non-coding portion of the genome. These findings have informed biology of lipoproteins and elucidated the causal roles of lipid levels on cardiovascular disease⁸⁻¹¹. Here, we build on these previous efforts to: 1) perform an exome-wide association screen for plasma lipids in >300,000 individuals; 2) evaluate discovered alleles experimentally in mouse models; and 3) test the inter-relationship of mapped lipid variants with four diseases – coronary artery disease (CAD), age-related macular degeneration (AMD), fatty liver, and type 2 diabetes.

We tested the association of genotypes from the HumanExome BeadChip (i.e., exome array) with lipid levels in each of 73 studies encompassing >300,000 participants (Supplementary Material, Tables S1-S3) across several ancestries (European, African, South Asian, Hispanic, and other). The maximal sample sizes from each ancestry were 237,050 for European, 16,935 for African, 37,613 for South Asian, and 5,082 for Hispanic or other. A companion manuscript describes results for 47,532 East Asian participants¹². A total of 242,289 variants were analyzed after quality control (Supplementary Methods). About one-third of the genotyped variants are non-synonymous with minor allele frequency (MAF) < 0.1% (Table S4).

Single-variant association statistics and inter-marker linkage disequilibrium information summarized across 1 megabase sliding windows were generated from each cohort using RAREMETALWORKER or RVTESTS^{13,14} software. Meta-analyses of genetic associations were performed using the R-package rareMETALS (version 6.0). In addition to testing each variant individually for association, we also performed gene-level association tests, which aggregate multiple rare coding variants in each of 16,798 protein-coding genes. Genomic control values for meta-analysis results were between 1.09 and 1.14 for all four lipid traits (Fig. S1), suggesting that population structure in our analysis is well-controlled^{3,15}. Genomic control values

scale with sample size¹⁶ and as such, our genomic control values are in line with previous reports including similar sample sizes¹⁵.

In single-variant association analyses, we identified 1,445 variants associated at $P < 2.1 \times 10^{-7}$ (Bonferroni correction of 242,289 variants analyzed) (Fig. S2; full association results are available at <http://csg.sph.umich.edu/abecasis/public/lipids2017/>). Of these, 75 were ‘novel’ (i.e., located in previously unreported loci): 35 of these were protein-altering variants and 40 were non-coding variants (**Table 1**, Table S5 & S6). These 75 new loci are each located at least 1 megabase from previously reported genome-wide association study (GWAS) signals (Table S7). The lead variant was common (MAF > 5%) in 61 of these 75 loci. When analyzed separately by ancestry, European ancestry participants provided the most significant associations for the 75 novel variants, except for two noncoding variants associated with LDL (rs201148465 and rs147032017) which were driven by the South Asian participants (Table S8). Gene-level association analyses revealed an additional five genes where the signal was driven by multiple rare variants ($P < 4.2 \times 10^{-7}$, Bonferroni correction threshold for performing 5 tests on ~20,000 genes, Table S9).

We tested the new single variant association findings for replication in an independent set of up to 286,268 participants from three studies – Nord-Trøndelag Health Study¹⁷, (HUNT; max n = 62,168), Michigan Genomics Initiative (MGI; www.michigangenomics.org; max n = 6,411) and the Million Veteran Program¹⁸ (MVP; max n = 218,117). Of the novel primary trait associations, 73/73 associations were directionally consistent (Table S10); two SNPs were not available for replication (rs201148465, rs75862065). Furthermore, we were able to replicate the associations of 66/73 (90%) at an alpha of 0.05.

At any given genetic locus, multiple coding and non-coding variants may contribute to plasma lipid levels in an independent fashion. We quantified this phenomenon by iteratively performing association analyses conditional on the top variants at all lipid loci (175 previously identified and 75 newly identified). With this procedure, we identified 444 variants independently associated with one or more of the four lipid traits in 250 loci (Fig. S3; Table S11

and Table S12). Collectively, these 444 variants explain 9%-12% of the overall phenotypic variance for plasma lipid levels; previously mapped variants explain about 5%-9% of the variance in the current dataset.

The identification of lipid-associated coding variants using the exome array may help refine association signals at previously identified GWAS loci. We were able to evaluate this possibility in 131 of the 175 previously reported GWAS loci where the index variant or a proxy was available on the exome array, and associated with lipids levels with $P < 2.1 \times 10^{-7}$ (Table S13 and Table S14). For example, rs11136341, an intronic SNP close to the *PLEC* gene reported to be associated with LDL-C, was the original lead SNP in its GWAS locus ($P = 2 \times 10^{-13}$). In the current study, a protein-altering variant in *PARP10* is the top variant in the same locus (rs11136343; Leu395Pro; $P = 7 \times 10^{-26}$). After conditioning on *PARP10* Leu395Pro, the evidence for rs11136341 diminished ($P = 0.02$); in contrast, *PARP10* Leu395Pro remained significant ($P = 9 \times 10^{-13}$) after conditioning on rs11136341. *PARP10* has been shown to affect the hepatic secretion of apolipoprotein B (apoB) in human hepatocytes¹⁹; these results prioritize *PARP10* as a causal gene at this locus.

Experimental analysis of discovered mutations in model systems is a powerful approach to validate human association findings. This approach holds particular appeal in following up the results of our exome array analysis, in which hundreds of protein-altering variants were robustly associated with circulating lipids. We prioritized two coding mutations for experimental analysis: *JAK2* (Janus Kinase 2) p.Val617Phe and *A1CF* (APOBEC1 complementation factor) p.Gly398Ser.

JAK2 p.Val617Phe is a recurrent somatic mutation arising in hematopoietic stem cells which can lead to myeloproliferative disorders or clonal hematopoiesis of indeterminate potential²⁰⁻²⁴. We recently showed that carriage of p.Val617Phe increases with age and confers higher risk for CAD²⁵. Surprisingly, the 617Phe allele which increases risk for CAD is associated with *lower* LDL-C. Mice knocked in for *Jak2* p.Val617Phe were created as reported previously²⁶. Hypercholesterolemia-prone mice that were engrafted with bone marrow obtained

from *Jak2* p.Val617Phe transgenic mice displayed lower total cholesterol than did mice that had received control bone marrow (Fig. S4). These data are consistent with our human genetic observations. The mechanism by which *JAK2* p.Val617Phe leads to lower plasma TC/LDL-C but higher risk for CAD requires further study.

Another strong new association to emerge from genetic analyses was between *A1CF* p.Gly398Ser and TG [0.10-standard deviation (SD) increase in TG per copy of alternate allele, $P = 4 \times 10^{-11}$]; this rare variant (MAF = 0.7%) was also associated with increased circulating TC ($P = 4 \times 10^{-7}$) and nominally associated with increased risk of CAD (OR 1.12; $P = 0.02$). *A1CF* encodes APOBEC1 complementation factor, an RNA-binding protein which facilitates the RNA-editing action of APOBEC1 on the *APOB* transcript^{27,28}. We performed CRISPR-Cas9 deletion, rescue, and knock-in experiments to assess whether *A1CF* p.Gly398Ser is a causal mutation that alters TG metabolism.

CRISPR-Cas9-induced deletion of *A1CF* led to 72% and 65% reduction in secreted APOB100 compared to control cells in Huh7 and HepG2 human hepatoma cells, respectively (**Fig. 1A-1C**; Fig. S5). These findings are consistent with previous studies in rat primary hepatocytes that also showed significantly decreased apoB secretion after RNAi-based depletion of *A1CF*²⁹. Additionally, cellular APOB100 levels were significantly reduced in *A1CF*-deficient cells (Fig. S5B and S5C). A subsequent “rescue” experiment involving overexpression of wild-type or p.Gly398Ser *A1CF* in Huh7 cells with or without endogenous *A1CF* expression confirmed that significantly more APOB100 secretion was noted in cell lines expressing *A1CF* p.Gly398Ser (**Fig. 1D**).

We sought to further validate the *A1CF* gene and the p.Gly398Ser variant through the use of CRISPR-Cas9 to generate knock-in mice. Using a guide RNA targeting *A1cf* exon 9, the site of the codon for p.Gly398, and a 162-nucleotide single-strand DNA oligonucleotide repair template containing the p.Gly398Ser variant as well as extra synonymous changes to prevent re-cleavage by CRISPR-Cas9, we generated mice of the C57BL/6J inbred background with an *A1cf* Gly398Ser allele (hereafter referred to as KI) (Fig. S6A, S6B). We bred the KI allele to

homozygosity and found that KI/KI mice were viable and healthy. We compared wild-type and KI/KI colony mates (n = 9, 8) with respect to TG levels (Fig. S6C). We found that KI/KI mice had 46% increased TG compared to wild-type mice ($P = 0.05$). In sum, these results indicate that *AICF* is a causal gene for TG in humans and that the p.Gly398Ser variant is a causal mutation, with possible relevance to CAD.

Next, we used the 444 identified DNA sequence variants to address four questions relevant to clinical medicine, focusing on one question for each of the four traits: TC, HDL-C, LDL-C, and TG. First, a rare null mutation in the beta-globin gene (*HBB*; c.92+1G>A, rs33971440) associated with lower total cholesterol (Table S15), raising the question of the relationship between beta-thalassemia and risk for CAD. Approximately 80 to 90 million individuals worldwide are estimated to carry a heterozygous loss-of-function *HBB* mutation, termed beta-thalassemia trait³⁰. Observational epidemiologic studies several decades ago showed that beta-thalassemia trait associates with lower blood cholesterol level^{31,32}. We find that *HBB* c.92+1G>A is not only associated with lower hematocrit but also with a 17 mg/dl decrease in LDL-C (-17 CI -23, -11; $P = 2.7 \times 10^{-8}$) and a 21 mg/dl decrease in TC (-21 CI -27, -14; $P = 8.9 \times 10^{-11}$) (Fig. S7). In an analysis of 31,156 CAD cases and 65,787 controls, carriers of loss-of-function variants in *HBB* were protected against CAD (odds ratio for CAD, 0.70; 95% CI 0.54, 0.90; $P = 0.005$, Fig. S8).

Second, DNA sequence variants in the *CETP* gene which associate with higher HDL-C also correlate with higher risk for age-related macular degeneration (AMD), a leading cause of blindness around the world³³⁻³⁷; here, we ask if *any way* of increasing *plasma* HDL-C will predictably lead to increased AMD risk. Across 168 independent HDL-C variants at 109 distinct loci with MAF > 1%, we tested the association of each independent HDL-C variant with AMD risk. The effect size of variant on HDL-C was positively correlated with its effect on AMD risk (correlation in effect sizes, $r = 0.41$, $P = 4.4 \times 10^{-8}$; Table S16, Fig. S9). However, this effect was driven by the 10 independent HDL-C associated variants in the *CETP* gene (heterogeneity across the different HDL-C-raising mechanisms ($\tau^2 = 0.91$, $P_{\text{het}} = 1.8 \times 10^{-15}$) (Table S17). A one-SD

increase in HDL-C via a genetic risk score of the 10 *CETP* variants robustly associated with increased AMD risk (OR = 2.01, 95% CI = 1.88-2.14, 16,144 AMD cases, 17,832 controls, $P = 3.4 \times 10^{-97}$). When these 10 *CETP* variants were removed, there was no longer a relationship between genetically-altered HDL-C and AMD risk ($P = 0.17$). These genetic results suggest that outside of the *CETP* locus, there is not a predictable relationship between plasma HDL-C and risk for AMD.

Third, will lowering LDL-C with lipid-modifying medicines always increase risk for type 2 diabetes? This question is motivated by the fact that in randomized controlled trials, statin therapy increases risk for type 2 diabetes^{26,27} and recent reports of *PCSK9* variants associating with higher risk for type 2 diabetes³⁸⁻⁴⁰. We confirmed the association of *PCSK9* p.Arg46Leu (R46L) with risk for type 2 diabetes among 222,877 participants (Table S18). We found that the 46Leu allele associated with lower LDL-C confers a 13% increased risk for type 2 diabetes (OR 1.13; 95% CI 1.06 – 1.20; $P = 6.96 \times 10^{-5}$) (Fig. S10). In addition, across 113 independent LDL-C variants at 90 distinct loci, we compared each variant's effect on LDL-C with its subsequent effect on risk for type 2 diabetes. Across the 113 variants, there is a weak inverse correlation between a variant's effects on LDL-C and type 2 diabetes ($r = -0.21$, $p = 0.025$); however, there is evidence for heterogeneity in this relationship ($\tau^2 = 0.50$, $P_{\text{het}} = 2.5 \times 10^{-9}$). Five LDL-C lowering genetic mechanisms had the most compelling evidence for association with higher risk for type 2 diabetes (*TM6SF2* p.Glu167Lys, *APOE* chr19:4510002, *HNF4A* p.Thr136Ile, *PNPLA3* p.Ile148Met, and *GCKR* p.Leu446Pro) ($P < 4.0 \times 10^{-4}$ for each, Bonferroni correction threshold for performing tests at 113 variants, Table S19; Fig. S11). These results suggest that only some ways of lowering LDL-C are likely to increase risk for type 2 diabetes.

Finally, two key processes – hepatic production and peripheral lipolysis – contribute to the blood level of TG. We asked how genes involved in hepatic production of TG-rich lipoproteins (*PNPLA3*, *TM6SF2*) differed from lipolysis pathway genes (*LPL*, *ANGPTL4*) in their impact on related metabolic traits - blood lipids, fatty liver, type 2 diabetes, and CAD (**Table 2**). The alternative alleles at *PNPLA3* p.Ile148Met, *TM6SF2* p.Glu167Lys, *LPL*

p.Ser474Ter, and *ANGPTL4* p.Glu40Lys all associated with lower blood triglycerides and reduced risk for CAD. However, the blood TG-lowering alleles at *PNPLA3* and *TM6SF6* led to more fatty liver and higher risk for type diabetes. In contrast, the blood triglyceride-lowering alleles at *LPL* and *ANGPTL4* were neutral with respect to fatty liver and led to lower risk for type 2 diabetes. We confirmed the *LPL* observation using a phenome-wide association study approach in the UK Biobank, a population-based cohort study (Table S20). In UK Biobank, a one-SD decrease in TG mediated by *LPL* variants reduced risks for *both* type 2 diabetes and CAD (**Fig. 2**).

In summary, combining large-scale human genetic analysis with experimental evidence, we demonstrate: (1) 444 independent coding and non-coding variants at 250 loci as associated with plasma lipids; (2) the use of mouse models and genome editing to pinpoint causal genes and protein-altering variants, as with example of *AICF* p.Gly398Ser; and (3) that *LPL* activation can be expected to lower triglycerides and reduce risks for *both* CAD and type 2 diabetes without increasing liver fat and thus be advantageous for patients with metabolic risk factors.

ONLINE METHODS

Study samples and phenotypes

Seventy-three studies contributed association results for exome chip genotypes and plasma lipid levels. The outcomes were fasting lipid values in mg/dL [TC, HDL-C, LDL-C, TG] from the baseline, or earlier exam with fasting measures. If a study only had non-fasting levels then it contributed only to the TC and HDL-C analyses. LDL-C and TG analyses were only performed on fasting lipid values. Lipid lowering therapy with statins was not routinely used prior to the publication of the 4S study in 1994 which demonstrated the clinical benefit of statin therapy. Therefore, for data collected before 1994, no lipid medication adjustment was applied. For data collected after 1994, we adjusted the TC values for individuals on lipid medication by replacing their total cholesterol values by $TC/0.8$; this adjustment estimates the effect of statins on TC values. No adjustment was made on HDL-C or TG. LDL-C was calculated using the Friedewald equation for those with $TG < 400$ mg/dl ($LDL-C = TC - HDL-C - (TG/5)$). If TC was modified as described above for medication use after 1994, then modified TC was used in this formula. If only measured LDL-C was available in a study, we used $LDL/0.7$ for those on lipid-lowering medication when data were collected after 1994. TG values were natural log transformed. For each phenotype, residuals were obtained after accounting for age, age^2 , sex, principal components (as needed by each study, up to four), and inverse normal transform residuals were created for analysis. For studies ascertained on CAD case/control status, the two groups were modeled as separate studies.

Genotyping and quality control

All studies assayed the Illumina or Affymetrix Human Exome array v1 or v1.1. Genotypes were determined from Zcall⁴¹ or joint calling⁴². Individual studies performed the following quality control: call rate, heterozygosity, gender discordance, GWAS discordance (if GWAS data available), fingerprint concordance, if available, and PCA outliers.

Association analyses

Each contributing cohort analyzed the ancestries within their cohorts separately and studies collected on case/control status analyzed cases separately from the controls. We performed both single variant and gene-level association tests. In the association analysis, we obtain residuals after controlling for sex, age, age² and up to 4 principal components as covariates. Studies that had related samples analyzed the association using linear mixed models with relatedness estimated from genome-wide SNPs or from pedigrees.

From each study, we collected single variant score statistics and their covariance matrix for variants in sliding windows across the genome. Summary association test statistics were generated using RAREMETALWORKER or RVTESTS. Using summary association statistics collected from each study, we performed meta-analysis of single variant association tests using the Mantel-Haenszel test and constructed burden, SKAT and variable threshold tests using the approach by Liu et al¹⁴. For burden and SKAT, we used minor allele frequency thresholds of 1% and 5% and for VT, we applied minor allele frequency threshold of 5%. In the SKAT test, variants are weighted according to their minor allele frequencies, using the beta kernel $\beta(1,25)$.

Using covariance matrices between single variant association statistics, we were also able to perform conditional association analyses centrally, which distinguishes genuine signals from “shadows” of known loci. Details of the methods can be found in Liu et al¹⁴.

We centrally performed quality control for the data. We aligned study reported reference and alternative alleles with alleles reported in the NHLBI Exome Sequencing Project⁴³ and remove mis-labelled variant sites that can be strand ambiguous. For variant sites in each study, we removed variants that had call rate < 0.9 or had Hardy Weinberg P values $< 1 \times 10^{-7}$. Finally, as additional checks, we visually inspected for each study the scatter plot of variant allele frequency against frequencies from ethnicity-matched populations in the 1000 Genomes Project⁴⁴, and made sure that the strand and allele labels were well calibrated between studies.

Single variant associations with $P < 2.1 \times 10^{-7}$ ($0.05/242,289$ variants analyzed) and gene-based associations with $P < 4.2 \times 10^{-07}$ ($0.05/[20,000 \text{ genes} * 6 \text{ tests}]$) were considered significant. Novel loci were defined as being not within 1 megabase of a known lipid GWAS SNP. Additionally, linkage disequilibrium information was used to determine independent SNPs where a locus extended beyond 1 megabase. All novel loci reported in this manuscript are > 1 megabase from any previously reported locus and independent ($r^2 < 0.2$ was required for variants within 3 megabases).

Sequential forward selection

To identify independently associated variants for each known and newly identified locus, we performed sequential forward selection. We initialized the set of independently associated variants (denoted by Φ), starting with the top association signal in the locus. For each iteration, conditioning on variants in Φ , we performed conditional association analyses for all remaining variants. If the top association signal after the conditional analysis remained significant, we added the top variant to the set Φ , and then repeated the conditional association analysis. If the top variant after the conditional analysis was no longer significant, we stopped and reported

variants in the set Φ as the final set of independent variants for that locus. We used the same single variant significance threshold ($P < 2.1 \times 10^{-7}$) to determine statistical significance with the sequential forward selection results (Fig. S3).

Annotation

Sequence variants were annotated according to refSeq version 1.9, using the SEQMINER software (version 5.7)⁴⁵. Transcript level annotations were obtained and prioritized. When multiple transcript level annotations were available, they were prioritized according to their functionality and deleteriousness. To implement gene-level association tests, the annotation with the highest priority was used (along with other filtering criteria such as minor allele frequencies) to determine the set of variants that are included.

Heritability and proportion of variance explained estimates

We estimated the proportion of variance explained by the set of 444 independently associated variants. The joint effects of variants in a locus were approximated by $\hat{\beta}_{JOINT} = \mathbf{V}_{META}^{-1} \vec{U}_{META}$, where \vec{U}_{META} is the single variant score statistics and \mathbf{V}_{META}^{-1} is the covariance matrix between them. The covariance between single variant genetic effects was approximated by the inverse of the variance-covariance matrix of score statistics, i.e. \mathbf{V}_{META}^{-1} . The phenotypic variance explained by the independently associated variants in a locus is given by $\hat{\beta}_{joint}^T \text{cov}(G) \hat{\beta}_{JOINT}$, where G is the genotypes of the analyzed variants.

Refinement of genome-wide association signals

We sought to quantify what proportion of GWAS loci might be due to a protein-altering variant and, therefore, directly identify a functional gene. We made the assumption that a protein-altering variant is the most likely causal variant for each region if it is the top signal, explains the signal, or is independent of the original signal. To identify putative functional coding variants accounting for the effects at known lipid loci, we performed reciprocal conditional analyses to control for the effects of known lipid GWAS or coding variants within 500kb, as this was the maximum distance for variants within the covariance matrix. Loci where coding variants are the most significant signals were considered as “coding as top”. Loci where the initial GWAS variants had conditional $P > 0.01$ were considered to be explained by the coding variants. Loci where the coding variants had conditional $P < 2.1 \times 10^{-7}$ were considered to be independent of the initial GWAS signals.

***JAK2* p.Val617Phe and plasma cholesterol in a mouse model**

Jak2 p.Val617Phe MxCre mice were created and reported previously²⁶. Bone marrow cells from the WT or *JAK2* p.Val617Phe MxCre mice, both treated with poly I:C, were transplanted into irradiated *Ldlr*^{-/-} recipients. After four weeks of recovery, the *Ldlr*^{-/-} recipient mice were fed a Western diet (TD88137, Harlan Teklad) for 8 weeks. Plasma was collected and 250 microliter of pooled plasma from 7 WT→*Ldlr*^{-/-} or 7 *Jak2* Val617Phe→*Ldlr*^{-/-} recipient was subjected to fast protein liquid chromatography on Sepharose CL-6B size exclusion column. Total cholesterol content in each fraction was assessed by Cholesterol E kit (Wako Diagnostics).

Validation of *AICF* with CRISPR-Cas9 in human cells

To knock out *AICF* in Huh7 and HepG2 human hepatoma cells, three CRISPRs (sgRNA sequences are as follows: 5'-GGATGCCCCCAACAAATAAT-3'; 5'-TTGGTTTTATCTGCAGCGCT-3' and 5'-ATGACTCTCATACTCCACGA-3') targeting exon 4 of the *AICF* gene were constructed by using the lentiviral vector lentiGuide-Puro. Packaged viruses were used to transduce the cells expressing Cas9 for 16 hours. Subsequently, cells were cultured in the presence of 5 µg/ml puromycin for five days before splitting for assays. Cells for APOB secretion assay were cultured for 18 hours in serum-free medium, then the amount of APOB100 in medium was measured using an ELISA kit (MABTECH) according to the manufacturer's instructions.

In a rescue experiment, to avoid cutting of the *AICF* coding region on the recombinant plasmids by previously designed exon-targeting CRISPRs, four new CRISPRs targeting introns flanking exon 4 were applied to deplete endogenous *AICF*. The sequences for those sgRNAs are as follows: 5'-GGTTGAAAATGTATGGCTGT-3'; 5'-TGATTGATGAGAATCCTTGT-3'; 5'-TTTTCTTTGCGGAGGCAGGC-3' and 5'-GCAGAGAAGTCTCATCTTTG-3'. The *AICF* p.Gly398Ser variant was generated by using overlapping PCR and confirmed by Sanger sequencing. Both wild-type and the *AICF* p.Gly398Ser variant were constructed into lentiviral plasmids, respectively. After transduction, cells were cultured for 48 hours in the presence of 100 ng/ml doxycycline to induce recombinant expression of *AICF* or p.Gly398Ser variant before performing different assays.

***Aicf* p.Gly390Ser knock-in mice**

All procedures used for animal studies were approved by Harvard University's Faculty of Arts and Sciences Institutional Animal Care and Use Committee and were consistent with local, state,

and federal regulations as applicable. Knock-in mice were generated using a guide RNA designed to target the orthologous site of the *AICF* p.Gly390Ser variant. In vitro transcribed Cas9 mRNA (100 ng/μL; TriLink BioTechnologies) and guide RNA (50 ng/μL) were co-injected with 100 ng/μL single-strand DNA oligonucleotide (Integrated DNA Technologies): (5'-GTTTGATTTCAGTTTAATGCCTTCTTACATTCCTAGAAATTTACATGAATGTCCCTGTA GGGGCTGCGGG GGTGAGAGGTCTGGGCAGCCGTGGCTATTTGGCATAACACAGGCCTGGGTCGAGGAT ACCACGTCAAAGGAGACAAGAGAGAAGACAAACTCTA-3') into the cytoplasm of fertilized oocytes from C57BL/6J mice. Genomic DNA samples from founder mice were screened for knock-in mutations by PCR and confirmed by Sanger sequencing. Positive mice were bred with C57BL/6J mice to generate wild-type and homozygous knock-in mice. Male colony mates at 12 weeks of age were used for lipid measurements. Blood samples were collected from the lateral tail vein following an overnight fast. Plasma triglyceride levels were measured using Infinity Triglycerides Reagent (Thermo Fisher) according to the manufacturers' instructions.

Intersection of lipid association signals with AMD, CAD, and T2D

To estimate the association of loss-of-function variants in *HBB* with cholesterol levels, participants from the following two consortia were studied: the Global Lipids Genetics Consortium and the Myocardial Infarction Genetics Consortium (MIGen, 27,939 participants in 12 cohorts). A rare loss-of-function variant in *HBB* (c.92+1G>A, rs33971440) was genotyped in participants from the Global Lipids Genetics Consortium Exome consortium. This variant was pooled with sequence data for the *HBB* gene in MIGen, available in 19,434 participants with

blood cholesterol measurements. The association of loss-of-function variants with cholesterol was estimated using linear regression with adjustment for age, sex and up to five principal components of ancestry. Estimates from genotyped and sequence data were pooled using inverse variance weighted fixed effects meta-analysis.

To estimate the association of loss-of-function variants in *HBB* with CAD, participants from the following two consortia were studied: the CARDIoGRAM Exome Consortium (69,087 participants from 20 studies) and MiGen (12384 CAD cases and 15547 controls from 12 studies). 69,086 individuals who were genotyped for the c.92+1G>A variant in CARDIoGRAM Exome were pooled with sequence data for *HBB* from 27,931 individuals in MiGen. The association of loss-of-function variants with CAD was estimated using logistic regression with adjustment for age, sex and up to five principal components of ancestry. Estimates were pooled using inverse variance weighted fixed effects meta-analysis. To estimate the association of loss of function variants in *HBB* with hemoglobin and hematocrit levels, estimates from an exome chip analysis of red blood cell traits (24 814 individuals) were used⁸.

For 168 variants independently and significantly associated with HDL-C and a MAF > 1%, we looked up the association evidence in 16,144 age-related macular degeneration cases and 17,832 controls with exome chip genotypes⁴⁶.

For 132 independently and significantly associated LDL-C variants and MAF > 1%, we looked up the association evidence in: (1) up to 120,575 individuals with and without CAD and exome chip genotypes (42,335 cases and 78,240 controls)⁴⁷; and (2) up to 69,870 individuals with and without type 2 diabetes. Only 113 of the 132 LDL variants were available in the type 2 diabetes results. We used a Bonferroni correction for 132 variants to determine significance of the results ($\alpha = 4.0 \times 10^{-4}$).

Association of *PCSK9* R46L with type 2 diabetes

For evaluating the association of *PCSK9* R46L with risk of type 2 diabetes, we considered a total of 42,011 type 2 diabetes cases and 180,834 controls from 30 studies from populations of European ancestry (Table S17). The variant was directly genotyped in all studies using the Metabochip or the Exome array. Sample and variant quality control was performed within each study as described previously⁴⁸⁻⁵¹. Within each study, the variant was tested for association with type 2 diabetes under an additive model after adjustment for study-specific covariates, including principal components to adjust for population structure. Association summary statistics for the variant for each study was corrected for residual population structure using the genomic control inflation factor as described previously⁴⁸⁻⁵⁰. We then combined association summary statistics for the variant across studies via fixed-effects inverse-variance weighted meta-analysis.

TG variants, lipids, fatty liver, type 2 diabetes, and CAD

Exome chip results for four variants (*LPL* p.Ser474Ter [rs328], *ANGPTL4* p.Glu40Lys [rs116843064], *PNPLA3* p.Ile148Met [rs738409], and *TM6SF2* p.Glu167Lys [rs58542926]) were obtained from the following sources:

1. lipids: current analysis
2. fatty liver: Between 2002 and 2005, 1,400 individuals from the Framingham Offspring Study and 2,011 individuals from third generation underwent multi-detector computed tomograms (CT) on which we evaluated liver attenuation as previously described⁵². We tested the association of TG variants with CT liver fat after inverse normal transformation. Covariates in the regression models included age, age², and gender. A similar analysis was conducted in 3,293 participants of

European ancestry from BioImage study⁵³. Association results for liver attenuation from the Framingham and BioImage studies were combined through fixed-effects inverse-variance weighted meta-analysis (Table S20).

3. type 2 diabetes: ExTexT2D Consortium⁵⁴

4. CAD: published results from the Myocardial Infarction Genetics and CARDIoGRAM Exome Consortia study⁴⁷.

Fig. 1: *A1CF* p.Gly398Ser mutant leads to increased APOB100 secretion. **A**, Western blot showing the depletion of endogenous A1CF levels via CRISPR/Cas9 system in both Huh7 and HepG2 cells. **B and C**, Lack of A1CF leads to reduced APOB100 secretion in Huh7 (B) and HepG2 (C) human hepatoma cells. **D**, Recombinantly overexpressed A1CF p.Gly398Ser variant led to significantly increased APOB100 secretion compared to A1CF or GFP control in both Huh7 wild-type and A1CF knockout cells (labeled as A1CF KO), respectively. The error bars show SD from experiments with biological replicates, N=6 for B, C and D. Statistically significant differences are marked (* $p < 0.05$, ** $p < 0.01$).

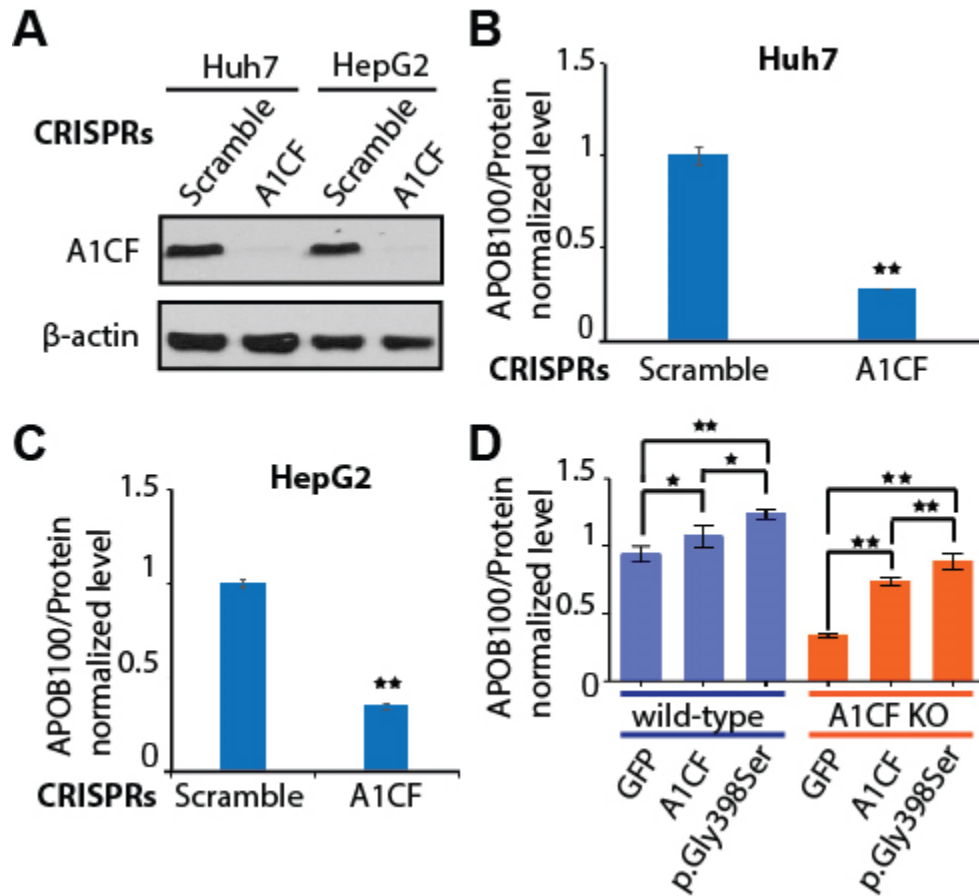


Fig. 2: Association of genetically-lowered triglycerides by *LPL* variants with a range of phenotypes. Estimates were derived in UK Biobank using logistic regression, adjusting for age, sex, ten principal components of ancestry and an indicator variable for array type. Effect estimates are for a 1 standard deviation lower plasma triglycerides. Definitions for all outcomes are provided in Table S16.

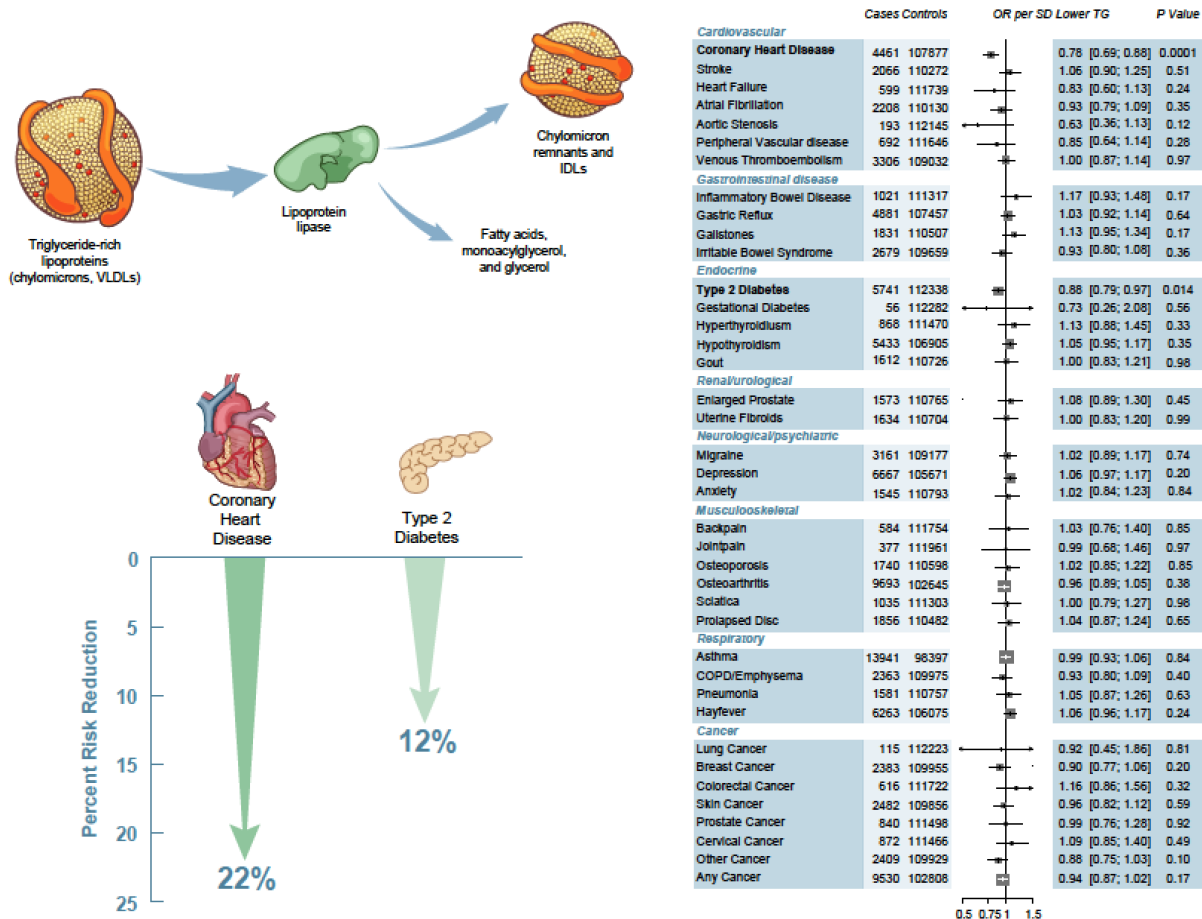


Table 1: Protein-altering variants at novel loci associated with lipid levels									
Chromosome: position (hg19)	rs ID	Alleles (reference/ alternative)	Gene	Protein change	N	Frequency alternative allele	Trait	P value	Beta±SE
Total Cholesterol									
2:101627925	rs1062062	C/T	<i>TBC1D8</i>	p.Gly954Arg	292898	0.12	TC	1×10 ⁻⁷	-0.021±0.0040
4:69343287	rs976002	A/G	<i>TMPRSS11E</i>	p.Tyr303Cys	293961	0.23	TC LDL-C	5×10 ⁻²⁰ 3×10 ⁻¹²	0.029±0.0031, 0.023±0.0033
4:155489608	rs6054	C/T	<i>FGB</i>	p.Pro206Leu	307997	0.0038	TC TG	5×10 ⁻¹² 3×10 ⁻¹¹	0.14±0.021, 0.14±0.021
9:5073770	rs77375493	G/T	<i>JAK2</i>	p.Val617Phe	188412	0.0011	TC LDL-C	1×10 ⁻¹¹ 2×10 ⁻⁹	-0.32±0.047, -0.30±0.049
9:117166246	rs2274159	A/G	<i>DFNB31</i>	p.Val400Ala	319677	0.48	TC	2×10 ⁻⁷	0.013±0.0026
17:8216468	rs871841	T/C	<i>ARHGEF15</i>	p.Leu277Pro	298725	0.52	TC	2×10 ⁻⁸	0.015±0.0026
19:18304700	rs874628	A/G	<i>MPV17L2</i>	p.Met72Val	319677	0.26	TC	2×10 ⁻⁷	0.015±0.0029
LDL Cholesterol									
1:155106227	rs4745	A/T	<i>EFNA1</i>	p.Asp137Val	291361	0.49	LDL-C	5×10 ⁻⁸	-0.015±0.0027
4:187120211	rs13146272	C/A	<i>CYP4V2</i>	p.Gln259Lys	295826	0.62	LDL-C	1×10 ⁻⁷	-0.015±0.0027
5:176520243	rs351855	G/A	<i>FGFR4</i>	p.Gly388Arg	233058	0.29	LDL-C	4×10 ⁻⁸	-0.018±0.0033
9:139368953	rs3812594	G/A	<i>SEC16A</i>	p.Arg1039Cys	293723	0.24	LDL-C	2×10 ⁻⁸	-0.018±0.0031
10:118397971	rs10885997	A/G	<i>PNLIPRP2</i>	p.Gln387Arg	258146	0.41	LDL-C	9×10 ⁻⁸	0.015±0.0029
10:124610027	rs1891110	G/A	<i>FAM24B</i>	p.Pro2Leu	295826	0.55	LDL-C TC	8×10 ⁻¹⁵ 2×10 ⁻¹³	0.021±0.0026, 0.019±0.0025
12:72179446	rs61754230	C/T	<i>RAB21</i>	p.Ser224Phe	292762	0.015	LDL-C	1×10 ⁻⁷	0.057±0.011
14:94844947	rs28929474	C/T	<i>SERPINA1</i>	p.Glu366Lys	290263	0.015	LDL-C TC	4×10 ⁻¹⁴ 6×10 ⁻¹⁴	0.081±0.011, 0.078±0.010
17:26694861	rs704	G/A	<i>VTN</i>	p.Thr400Met	295826	0.49	LDL-C TC	6×10 ⁻¹⁶ 1×10 ⁻⁸	0.021±0.0026, 0.015±0.0025
19:42584958	rs201596848	C/T	<i>ZNF574</i>	p.Arg734Cys	273744	0.0014	LDL-C	5×10 ⁻¹²	-0.255±0.037
Triglycerides									
2:202122995	rs3769823	A/G	<i>CASP8</i>	p.Lys14Arg	295956	0.69	TG	1×10 ⁻⁹	0.017±0.0028
5:131008194	rs26008	T/C	<i>FNIP1</i>	p.Gln620Arg	305699	0.92	TG	5×10 ⁻⁹	-0.028±0.0048
10:52573772	rs41274050	C/T	<i>A1CF</i>	p.Gly398Ser	299984	0.0072	TG TC	4×10 ⁻¹¹ 1×10 ⁻⁷	0.10±0.015, 0.08±0.015
13:45970147	rs138358301	A/G	<i>SLC25A30</i>	p.Phe280Leu	301087	0.0035	TG	3×10 ⁻¹¹	0.15±0.022
15:40751555	rs3803357	C/A	<i>BAHD1</i>	p.Gln298Lys	305699	0.55	TG	1×10 ⁻¹⁰	-0.017±0.0026
17:17409560	rs7946	C/T	<i>PEMT</i>	p.Val212Met	304420	0.67	TG	1×10 ⁻⁸	-0.016±0.0029
20:56140439	rs41302559	G/A	<i>PCK1</i>	p.Arg483Gln	299984	0.0021	TG	9×10 ⁻⁸	-0.154±0.029
22:17625915	rs35665085	G/A	<i>CECR5</i>	p.Thr149Met	302582	0.050	TG	5×10 ⁻⁸	0.032±0.0059
HDL Cholesterol									
2:272203	rs11553746	C/T	<i>ACPI</i>	p.Thr95Ile	313148	0.33	HDL-C	5×10 ⁻⁸	0.015±0.0027
2:54482553	rs17189743	G/A	<i>TSPYL6</i>	p.Arg246Cys	314415	0.029	HDL-C	2×10 ⁻⁷	0.040±0.0076
2:179309165	rs75862065	G/A	<i>PRKRA</i>	p.Pro116Leu	105490	0.29	HDL-C	2×10 ⁻⁷	0.026±0.0050
3:48229366	rs146179438	C/A	<i>CDC25A</i>	p.Gln25His	288306	0.020	HDL-C	3×10 ⁻¹¹	-0.063±0.0095
5:176637576	rs28932178	T/C	<i>NSD1</i>	p.Ser457Pro	310567	0.17	HDL-C	8×10 ⁻⁹	0.020±0.0035
11:64031241	rs35169799	C/T	<i>PLCB3</i>	p.Ser778Leu	314415	0.060	HDL-C TG	4×10 ⁻¹³ 3×10 ⁻¹²	-0.039±0.0054, 0.038±0.0055
11:68703959	rs622082	A/G	<i>IGHMBP2</i>	p.Thr671Ala	316391	0.31	HDL-C	6×10 ⁻¹⁰	-0.017±0.0028

16:4755108	rs78074706	G/A	<i>ANKS3</i>	p.Arg286Trp	315298	0.022	HDL-C	1×10^{-9}	-0.053±0.0087
16:69385641	rs76116020	A/G	<i>TMED6</i>	p.Phe6Leu	310822	0.033	HDL-C	7×10^{-9}	-0.041±0.0071
17:40257163	rs2074158	T/C	<i>DHX58</i>	p.Gln425Arg	244331	0.19	HDL-C	1×10^{-7}	-0.020±0.0038

Table 2. Impact of genes involved in hepatic production of triglyceride-rich lipoproteins (*PNPLA3*, *TM6SF2*) versus lipolysis pathway genes (*LPL*, *ANGPTL4*) on related metabolic traits - blood lipids, fatty liver, type 2 diabetes, and coronary artery disease.

Gene	<i>LPL</i>	<i>ANGPTL4</i>	<i>PNPLA3</i>	<i>TM6SF2</i>
Variant	p.Ser474Ter	p.Glu40Lys	p.Ile148Met	p.Glu167Lys
Effect Allele	X	K	Met	K
Frequency	10%	2%	23%	7%
Blood triglycerides				
Effect Direction	↓	↓	↓	↓
Beta	-0.18	-0.27	-0.018	-0.12
(SE)	(0.004)	(0.010)	(0.003)	(0.005)
P	$P < 1 \times 10^{-323}$	$P = 4 \times 10^{-175}$	$P = 4 \times 10^{-9}$	$P = 4 \times 10^{-125}$
Blood LDL cholesterol				
Effect Direction	-	↓	↓	↓
Beta	0.013	-0.004	-0.018	-0.103
(SE)	(0.004)	(0.010)	(0.003)	(0.005)
P	$P = 0.005$	$P = 0.70$	$P = 1 \times 10^{-8}$	$P = 7 \times 10^{-93}$
Fatty liver				
Effect Direction	-	-	↑	↑
Beta*	0.026	0.112	-0.247	-0.250
(SE)	(0.031)	(0.068)	(0.022)	(0.036)
P	$P = 0.41$	$P = 0.10$	$P = 2 \times 10^{-30}$	$P = 5 \times 10^{-12}$
Type 2 diabetes				
Effect Direction	↓	↓	↑	↑
Beta	-0.049	-0.097	0.039	0.07
(SE)	(0.01)	(0.040)	(0.005)	(0.01)
P	$P = 7 \times 10^{-9}$	$P = 1 \times 10^{-4}$	$P = 2 \times 10^{-10}$	$P = 5 \times 10^{-12}$
Coronary artery disease				
Effect Direction	↓	↓	↓	↓
Beta	-0.074	-0.160	-0.044	-0.047
(SE)	(0.014)	(0.026)	(0.008)	(0.013)
P	$P = 4 \times 10^{-7}$	$P = 2 \times 10^{-10}$	$P = 4 \times 10^{-8}$	$P = 3 \times 10^{-4}$

*A negative beta reflects liver attenuation on computed tomography which is indicative of *higher* liver fat

Association results for lipids are derived from present study

Association results for type 2 diabetes are from⁵⁴

Association results for coronary artery disease are from⁴⁷

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